



International Journal of Pharmacy and Biological Sciences ISSN: 2321-3272 (Print), ISSN: 2230-7605 (Online) IJPBS™ | Volume 8 | Issue 3 | JUL-SEPT | 2018 | 579-592

Research Article | Biological Sciences | Open Access | MCI Approved| ज्ञान-विज्ञान विमुक्तये |UGC Approved Journal |

COMPARATIVE MORPHOLOGICAL AND PHYTOCHEMICAL ANALYSES OF THREE VARIANTS OF *CISSUS QUADRANGULARIS* IN TAMIL NADU

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ABSTRACT

To study Morpho-parameters and the Phyto-constituents of Cissus quadrangularis (L.). variant I, II and III to provide its chemical constituents and find out the morphological variations between the three variants available in Tamil Nadu. The dried and powered stem materials (20 g) were extracted successively with 200 mL of methanol. Phytochemical analyses were carried out by using GC-MS and FT-IR. GC-MS identify the chemical compounds and FT-IR used to identify the functional group of Compounds. Based on GC-MS methanolic stem extract of C. quadrangularis three variants showed similar major compounds were identified. Such as 3- o- methyl- d- glucose, D-Allose, Phytol and 9, 12, 15 – octadecatrienoic acid. The FT-IR analysis showed similar functional group of compounds. The morphological character of C. quadrangularis stomed various type of stems variation. Based on the morphological characters of C. quadrangularis stem shape differs in the three variants studied. Based on the GC-MS and FT-IR analyses similar phyto- constituent related to functional groups were present in all the three variants studied. The all the three variants of C. quadrangularis morphologically different in nature but the phytochemical study showed no major differents due to environmental adaptation of these plants were different from morphological character.

KEY WORDS

C. quadrangularis, FT-IR, GC-MS, Morphology, Phyto-constituents

INTRODUCTION

The world is rich with natural and unique medicinal and aromatic plants. Medicinal plants are now getting more devotion than ever because they have potential of numerous benefits to society or indeed to all mankind, especially in the line of medicine and pharmacological. The medicinal value of these plants lies in bioactive phytochemical constituents that produce definite physiological action on the human body [1]. The medicinal plants are valuable for curing of human diseases because of the presence of phytochemical constituents [2]. Phytochemicals are constituents commonly existing in plants. They give to the dye (colour), flavour and smell of plants. In addition, they form part of a plant's natural protection mechanism against diseases. Their medicinal values to human health and disease prevention have been reported [3]. Intraspecific variation of chemical compounds is common in many plant species and often shows clear geographical patterns that may reflect environmental differences within the range of a species. Hence, chemical races are often defined and provide taxonomists with a powerful tool [4]. The genus Cissus with 350 species is distributed in the world. In India, the genus is signified by approximately 13 species [5] and in Tamil Nadu state by approximately 11 species were occurred [6]. Cissus quadrangularis Linn. belongs to the family Vitaceae is a climber with stout fleshy quadrangular (four angled) stem found throughout the hotter parts of India [7]. Phytochemical studies of C. quadrangularis variant found numerous phytochemical



constituents such as ascorbic acid, carotene, anabolic steroidal substances, calcium, beta-sitosterol, alphaamyrin, alpha- amyrone, flavonoids, triterpenoids [8] and various secondary metabolites. Based on the stem Morphology the plant C. quadrangularis divided into three variants such as square-stemmed, roundstemmed and flat-stemmed are available. They are differentiated as variant I, II and III respectively [9]. C. quadrangularis variant I so called wild plant compare with other two variants. Pharmacological activities of C. quadrangularis have been reported in earlier [10-11]. In the present study to investigate the morphology and phytochemical variations of C. quadrangularis of three variants.

MATERIALS AND METHODS

Collection and preparation of plant material

The fresh stem plant material of C. quadrangularis variant I, II and III were collected from Chidambaram and Cuddalore. The herbarium specimen of the same were prepared and deposited in the Department of Botany, Annamalai University. The plants were washed completely in running tap water to remove soil particles and the plant parts were separated and shade dried. The shade dried plant parts were stored in air tight container for further analysis.

Morphological Characters of C. quadrangularis

Climbing herb, tendrils simple, opposite to the leaves, leaves simple or lobbed, sometimes 3 - foliate, dentate. Flowers bisexual, tetramerous, in umbellate cymes, opposite to the leaves, Calyx cup- shaped, obscurely 4-lobed. Fruit globose or obovoid fleshy berries, one seeded dark purple to black, seeds ellipsoid or pyriform. Were Observed morphological variations of C. quadrangularis Variant I, II and III.

Plant sample extraction

Phytochemical analyses were carried out by using Maceration methods [12]. Shade dried and powdered plant materials were successively extracted with methanol with gentle stirring for 72 hours. The extraction were passed through Whatmann No. 1 paper and collected.

Gas Chromatography- Mass Spectrum Analysis (GC-MS)

25 grams of the stem and leaves powder of C. guadrangularis variant I, II and III were soaked in 95 % methanol for 12 hours. The extracts were then filtered through Whatmann filter No. 41 along with 2 gm Sodium sulpate to remove the sediments and traces of water in the filtrate. Then the filtrate was concentrated by introducing bubbling nitrogen gas into the solution. The plant extract contains both polar and non-polar phytochemicals. 2 μ l of the plant methanolic extract filtrate was used for GC-MS analysis. GC-MS analysis of the methanolic extract of the plant samples taken for this study was performed by using a Perkin- Elmer GC clarus- 500 system comprising an AOC 20°C auto sampler and a Gas chromatograph intergraded to a mass spectrometer equipped with a Elite - 5MS fused capillary column. Helium gas was used as a carrier gas at a constant flow rate of 1 ml/min, and an injector where of 2 µl was employed. The inject temperature was maintained at 250°C, the ion source temperature was 200°C and oven temperature was programmed from 110°C, with an increase of 10°C/min to 200°C, then 5°C/min 280°C ending with a 9 min isothermal at 280°C. The solvent delay was 0 to 2 min, and the total GC/MS running time was 36 min. The relative percentage amount of each component was calculated by comparing its average peak area to the total areas. The Mass detector used in this analysis was turbo- Mass gold Perkin- Elmer and the software adopted to handle Mass spectra and chromatogram was a Turbo – Mass Ver.5.2. **FT-IR** analysis

The FT-IR analysis was done by utilizing Perkin Elmer Spectrum Version 10.03.09 framework, which was utilized to identify the practical gatherings of the compound. A little measure of concentrate was set specifically on the zinc solenoid piece and consistent weight. Information of infrared retentive, gathered over the wave number extended from 4000 cm⁻¹ to 400 cm⁻¹ utilizing spectra programming. Tests were keep running in triplicate and every one of them were embraced inside a day time span.

RESULTS AND DISCUSSION

Morphological variations of C. quadrangularis

Table 1. showed morphological characters of all the three variants of C. quadrangularis showed similar character of Root, Tendril, Flower, Fruit and Seed. The major difference in all three variants were observed in stem, C. quadrangularis Variant I having 4 angled stem, C. quadrangularis Variant II - cylinder shaped stem and C. quadrangularis Variant III were observed flat shaped stem. Minor different were observed in leaf character. C. quadrangularis Variant I and II having simple leaves



and variant III were observed palmate leaves. (Figure 1, 2 and 3)

S.No.	Character	C. quadrangularis Variant I	<i>C. quadrangularis</i> Variant II	C. quadrangularis Variant III
1	Habit	Climber	Climber	Climber
2	Stem	Quadrangular Stems from 1 - 12 meters (Avg,) long from a tuberous rootstock	Cylinder Stems from 1-13 meter (Avg,) long from a tuberous rootstock	Flat Stems from 12 meter (Avg,) long from a tuberous rootstock
3	Root	Tap-root 10-20 Cm (Avg,)	Tap-root 8-15 Cm (Avg,)	Tap-root 5-13 Cm (Avg,)
4	Leaf	Simple, alternate, 2-6 Cm (Avg,)	Simple, alternate 2-5 Cm (Avg,)	Alternate, palmate 2-6 Cm (Avg,)
5	Tendril	These simple tendrils are long, stout, slender and leaf-opposed. 8-20 Cm (Avg,)	These simple tendrils are long, stout, slender and leaf-opposed. 8-15 Cm (Avg,)	These simple tendrils are long, stout, slender and leaf- opposed. 8-20 Cm (Avg,)
6	Flower	Umbellate cymes yellowish-Green 0.8 Cm (Avg,)	Umbellate cymes yellowish-Green 0.8 Cm (Avg,)	Umbellate cymes yellowish-Green 0.8 Cm (Avg,)
7	Fruit	Globose, succulent berry, 0.6 to 1 Cm (Avg,) These fruits are green turning red when ripe.	Globose, succulent berry, 0.6 to 1 Cm (Avg,), These fruits are green turning red when ripe.	Globose, succulent berry, 0.6 to 1 cm (Avg,), These fruits are green turning red when ripe.
8	Seed	Small with thick Testa	Small with thick Testa	Small with thick Testa

Table 1. Morphological Character of *C. quadrangularis* variants



Figure 1: Morphology of C. quadrangularis variant I





Figure 2: Morphology of C. quadrangularis variant II



Figure 3: Morphology of *C. quadrangularis* variant III

GC-MS

The phytochemical constituents present in the stem methanolic extract of *C. quadrangularis* variant I stem showed fifty-four phytochemical constituents. Out of these fifty-four constituents five constituents were majorly present such as 3-O-Methyl D-glucose, D-Allose, 9, 12, 15-Octadecatrienoic acid, Phytol and Pentadecanoic Acid. Based on GC-MS spectrum confirmed the compounds with retention time 23.574, 19.237, 28.706, 25.037and 26.250 respectively (Figure 4). Apart from the above-mentioned compounds, the 3-

O-Methyl-D-glucose was containing the percentage of 51.85 and it was identified as active compound of the species *C. quadrangularis* variant I. The molecular formula of the compound was $C_7H_{14}O_6$ (Table 2). GC-MS analysis of Methanolic extract of *C. quadrangularis* variant II stem demonstrates nearness of fifty-six constituents. Out of these fifty-six constituents five constituents were majorly present such as Hexadeconic acid, Ergost – 5-en- 3- ol, D-Allose, Phytol and 3-O-Methyl D-glucose with their respective retention time such as 27.425, 39.178, 39.453, 40.368 and 41.122



respectively (Figure 5). Based on the spectrum percentage area, Phytol (18.23) was identified as major active compound and the molecular formula was $C_{20}H_{40}O$ (Table 3). Methanolic stem extract of *C. quadrangularis* variant III indicates thirty-four constituents. There are five major constituents were present likely Hexadeconic acid, D-Allose, 9, 12, 15-

Octadecatrienoic acid, Phytol and 3-O-Methyl Dglucose. Based on GC-MS spectrum confirmed the compounds with retention time 45.866, 46.005, 46.760, 47.285 and 47.853 respectively (Figure 6). The D- Allose (23.24) was identified as active compound of the *C. quadrangularis*. The molecular formula of the compound was $C_6H_{12}O_6$ (Table 4).

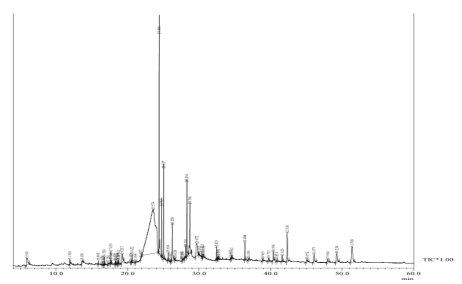


Figure 4: GC-MS Chromatogram of methanolic extract of C. quadrangularis stem variant I

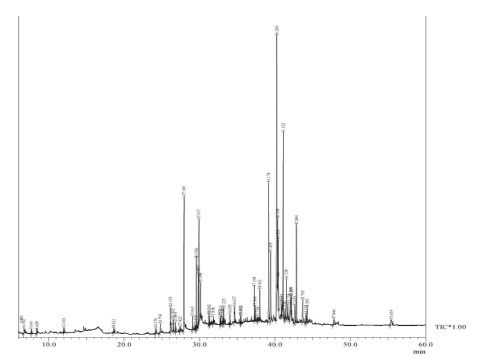


Figure 5: GC-MS Chromatogram of methanolic extract of C. quadrangularis Stem Variant II

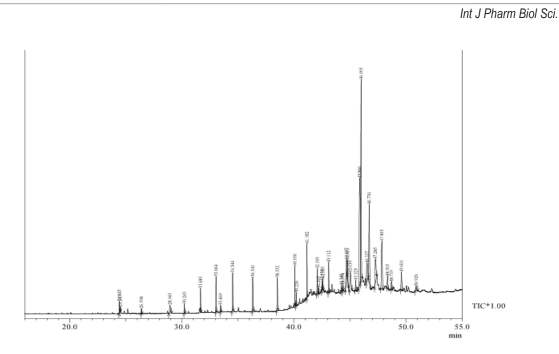


Figure 6: GC-MS Chromatogram of methanolic extract of C. quadrangularis Stem variant III

Peak	R. Time	Area%	Name of the compounds	Molecular Formula	Molecular weight
1	5.910	1.48	1H-Pyrrole	C4H5N	67
2	11.910	0.32	2,3-Dihydro-3,5-Dihydroxy-6-Methyl-4h- Pyran-One	$C_6H_8O_4$	144
3	13.630	0.31	2h-Pyran-2-One, 5,6-Dihydro-6-Pentyl-	$C_{10}H_{16}O_2$	168
4	15.895	0.12	2. Alpha., 7, 8-trimethylacenaphthylene	$C_{15}H_{24}$	204
5	16.439	0.04	2-(3-Isopropyl-4-Methyl-3-Penten-1-Ynyl)-2- Methycyclobutanone	C ₁₄ H ₂₀ O	204
6	16.663	0.04	3,6,6,7-Tetramethyl-3-Vinyl-2,3,3a,4,5,6- Hexahydro-1h-Indene	$C_{15}H_{24}$	204
7	16.753	0.25	Tricyclo[4.4.0.0(2,7)]Dec-3-Ene, 1,3-Di Methyl -8-(1-Methylethyl)	C15H24	204
8	17.281	0.04	Aromadenrene	$C_{15}H_{24}$	204
9	17.635	0.36	Trans (.beta.)-caryophyllene	C15H24	204
10	17.689	0.07	Isoledene	C15H24	204
11	18.27	0.11	1,4,8-Cycloundecatriene ,2,6,6,9-Tetra Methyl	C15H24	204
12	18.412	0.09	1H-Cycloprop [E]Azulene,	$C_{15}H_{24}$	204
13	18.634	0.10	6.AlphaCadina-4,9-Diene, (-)-	C15H24	204
14	18.762	0.20	Betaylangene	C15H24	204
15	19.237	9.28	D-Allose	$C_6H_{12}O_6$	180
16	20.443	0.03	1-Tridecanol	C13H28O	200
17	20.622	0.31	(-)-5-Oxatricyclo [8.2.0.0(4,6)] Dodecane,	$C_{15}H_{24}O$	220
18	21.064	0.05	Humulene Oxide	C ₁₅ H ₂₄ O	220
19	21.837	0.07	(1ar-(1aalpha, 5abeta, 9ar(*)))-5a,9,9- Trimethyloctahydrobenzo(D)Cycloprop(C)O xepin-2,4	C ₁₄ H ₂₀ O ₃	236
20	23.574	51.85	3-O-Methyl-d-glucose	C7H14O6	194
21	24.406	1.55	2,6,10-Trimethyl,14-Ethylene-14- Pentadecne	C ₂₀ H ₃₈	278
22	24.765	2.15	Neophytadiene	C ₂₀ H ₃₈	278

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23	25.037	3.41	Phytol	C ₂₀ H ₄₀ O	296
24	25.654	0.37	Hexadecanoic Acid,	$C_{17}H_{34}O_2$	270
25	25.993	0.05	Isophytol	C ₂₀ H ₄₀ O	296
26	26.250	3.39	Pentadecanoic Acid	$C_{15}H_{30}O_2$	242
27	26.628	0.08	14B-Pregnane	$C_{21}H_{36}$	288
28	27.597	0.0	Methyl 4-o-benzylalphal-rhamnopyra noside	$C_{14}H_{20}O_5$	268
29	27.690	0.04	1H-Indene, 3-Methyl-	$C_{10}H_{10}$	130
30	28.116	0.63	9,12,15-Octadecatrie noic Acid, (Z,Z,Z)-	$C_{19}H_{32}O_2$	292
31	28.294	2.72	3,7,11,15-Tetramethyl-, [R-[R*,R*-(E)]]	C ₂₀ H ₄₀ O	296
32	28.706	3.83	9,12,15-Octadecatrienoic Acid,	$C_{18}H_{30}O_2$	278
33	29.672	2.93	Cyclohexanol, 4-[(Tri Methylsilyl) Oxy] -, Cis-	C ₉ H ₂₀ O ₂	188
34	30.015	0.25	Decanoic Acid	$C_{10}H_{20}O_2$	172
35	30.422	0.20	Octanoic acid, 2-di methylaminoethyl ester	$C_{12}H_{25}NO_2$	215
36	30.630	0.33	5-Phenyl-Pentanoic Acid, Ethyl Ester	C17H24O5	308
37	32.42	0.42	Fumaric acid, 2-di methyl aminoethylnonyl ester	C ₁₇ H ₃₁ NO ₄	313
38	32.610	0.06	3-Cyclopentylpropionic Acid,	$C_{12}H_{23}NO_2$	213
39	32.715	0.13	2-Ethylbutyric Acid, Eicosyl Ester	C ₂₆ H ₅₂ O ₂	396
40	34.397	0.10	1-Penten-3-One, 4-Methyl-1-Phenyl-	$C_{12}H_{14}O$	174
41	34.562	0.20	Ethyl linolate	C ₁₉ H ₃₂ O ₂	292
42	36.408	0.79	Squalene	C ₃₀ H ₅₀	410
43	36.914	0.18	TriacontylPentafluoropropionate	C33H61F5O2	584
44	38.909	0.15	Chol-5-ene-3,24-diol	C ₂₄ H ₄₀ O ₂	360
45	39.715	0.30	Ergost-5-En-3-Ol, (3. Beta.)-	C ₂₈ H ₄₈ O	400
46	40.396	2.98	Beta-Sitosterol	C ₂₉ H ₅₀ O	414
47	40.854	0.10	5-chlorostigmastan-3-yl acetate	$C_{31}H_{53}C_1O_2$	492
48	41.626	0.42	Stigmast-5-En-3-Ol, (3. Beta.)-	C ₂₉ H ₅₀ O	414
49	42.318	2.70	Vitamin E	C ₂₉ H ₅₀ O ₂	430
50	4\$5.092	0.40	24- Epicampesterol	C ₂₈ H ₄₈ O	400
51	46.073	1.05	Stigmasta-5,22-Dien-3-ol	C ₂₉ H ₄₈ O	412
52	47.994	0.52	Beta-di hydrofucosterol	$C_{29}H_{50}O$	414
53	49.254	1.50	,14,14a,14b-Octadbctadecahyro-2h-Picen-3- One	C ₃₀ H ₄₈ O	424
54	51.398	0.89 100.00	D:B-Friedo-B':A'-Neogammacer-5-En-3-one	C ₃₀ H ₄₈ O	424

Table 3: GC-MS report of Methanolic extract of C. quadrangularis stem Variant II

Peak	R. Time	Area%	Name of the compounds	Molecular formula	Molecular weight
1	5.237	0.56	1-Butanamine, 2-methyl-N-(2 methylbutylidene)-	$C_{10}H_{21}N$	155
2	5.435	0.69	2,3-lupetidine	$C_{10}H_{21}N$	155
3	5.881	0.52	2-Pyrrolidinone	C ₄ H ₇ NO	85
4	6.692	0.51	Benzeneethanamine	$C_8H_{11}N$	121
5	7.690	0.14	2,3-Dihydro-3,5-Dihydroxy-6- Methyl-4H-Pyran-4- one	$C_6H_8O_4$	144
6	8.428	0.16	5-Methoxypyrrolidin-2-one	C ₅ H ₉ NO ₂	115
7	12.001	0.26	2-Piperidineacetic acid, Alpha- phenyl-,Methylester,Threo-	C14H19NO2	233
8	18.621	0.17	Dodecanoic acid	C12H24O2	200
9	24.176	0.23	4-((1E)-3-Hydroxy-1- propenyl)-2-methoxyphenol	C ₁₀ H ₁₂ O ₃	180
10	24.794	0.60	Pentadeconic acid	C15H30O2	242

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11	26.155	0.79	2,6,10-Trimethyl,14-ethylene-	C ₂₀ H ₃₈	278	
12	26.502	0.23	14-pentadecne Tetradecanoic acid	C14H28O2	228	
13	26.804	0.34	3,7,11,15-Tetramethyl-2-	C ₂₀ H ₄₀ O	296	
13	20.004	0.54	hexadecen-1-ol	C20H40U	290	
14	27.425	0.12	Hexadecanoic acid, methyl ester	C ₁₇ H ₃₄ O ₂	270	
15	27.945	8.00	Hexadecanoic acid	C ₁₆ H ₃₂ O ₂	256	
16	29.065	0.47	Heptadecanoic acid	$C_{16}H_{34}O_2$	270	
17	29.476	0.15	9,12,15-octadecatrienoic acid, Methyl ester	C19H32O2	292	
18	29.596	2.10	2-hexadecen-1-ol, 3,7,11,15- tetramethyl-, [R-[R*,R*-(E)]]-	$C_{20}H_{40}O$	296	
19	29.857	2.73	Cyclodecene	C ₁₀ H ₁₈	138	
20	29.917	1.11	7-Tetradecenal, (Z)-	C14H26O	210	
21	30.144	1.13	Octadecanoic acid	$C_{18}H_{36}O_2$	284	
	001211		3-Cyclopentylpropionic acid,	018.15002	20.	
22	31.282	0.26	2-	$C_{12}H_{23}NO_2$	213	
	51.202	0.20	dimethylaminoethyl ester	C1211231102	213	
23	31.339	0.12	N- Nonadecanol-1	C19H40O	284	
23	21.222	0.12	Cyclobutanecarboxylic acid,	C191140U	204	
24	31.818	0.21		$C_{16}H_{28}O_2$	252	
			undec-2-enyl ester			
25	22 757	0.40	Fumaric acid, 2-		242	
25	32.757	0.10	dimethylaminoethyl nonyl	C ₁₇ H ₃₁ NO ₄	313	
			ester			
26	32.814	0.22	Fumaric acid, 2-di methyl	C ₁₅ H ₂₇ NO ₄	285	
			amino ethyl heptyl ester			
27	33.082	0.22	n-Tetracosanol-1	C ₂₄ H ₅₀ O	354	
28	33.225	0.40	Hexadecanoic acid, 2-hydroxy- 1-(hydroxymethyl)ethyl ester	C19H38O4	330	
29	34.053	0.33	Cyclohexane, 1,1'- tetradecylidenebis-	C ₂₆ H ₅₀	362	
30	34.657	1.22	cis-9-Hexadecenal	C ₁₆ H ₃₀ O	238	
31	35.462	0.17	Heptacosyl acetate	C ₂₉ H ₅₈ O ₂	438	
32	35.513	0.18	Squalene	C ₃₀ H ₅₀	410	
33	37.298	1.08	.gammaTocopherol	C ₂₈ H ₄₈ O ₂	416	
55			Cholest-24-en-16-one,			
34	37.369	0.39	(5. alpha.,20.xi.)-	C ₂₇ H ₄₄ O	384	
25		0.20	1-heneicosyl formate	$C_{22}H_{44}O_2$	340	
35	37.718	0.36			0.10	
			Vitamin E			
36	38.002	3.73	-	C ₂₉ H ₅₀ O2	430	
			Vitamin E			
36	38.002	3.73	Vitamin E Ergost-5-en-3-ol, (3.	C ₂₉ H ₅₀ O2	430	
36 37	38.002 39.178	3.73 7.54	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)-	C ₂₉ H ₅₀ O2 C ₂₈ H ₄₈ O	430 400	
36 37 38	38.002 39.178 39.453	3.73 7.54 5.24	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose	C29H50O2 C28H48O C6H12O6	430 400 180	
36 37 38	38.002 39.178 39.453	3.73 7.54 5.24	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol	C29H50O2 C28H48O C6H12O6	430 400 180	
36 37 38 39	38.002 39.178 39.453 40.269	3.73 7.54 5.24 18.23	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol 2,4a,8,8-Tetramethyl decahydro	C ₂₉ H ₅₀ O2 C ₂₈ H ₄₈ O C ₆ H ₁₂ O ₆ C ₂₀ H ₄₀ O	430 400 180 296	
36 37 38 39	38.002 39.178 39.453 40.269	3.73 7.54 5.24 18.23 3.10	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol 2,4a,8,8-Tetramethyl decahydro cyclopropa[d]naphthalene	C ₂₉ H ₅₀ O2 C ₂₈ H ₄₈ O C ₆ H ₁₂ O ₆ C ₂₀ H ₄₀ O	430 400 180 296	
36 37 38 39 40	38.002 39.178 39.453 40.269 40.368	3.73 7.54 5.24 18.23	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol 2,4a,8,8-Tetramethyl decahydro cyclopropa[d]naphthalene Olean-12-en-3-one	C ₂₉ H ₅₀ O2 C ₂₈ H ₄₈ O C ₆ H ₁₂ O6 C ₂₀ H ₄₀ O C ₁₅ H ₂₆	430 400 180 296 206	
36 37 38 39 40 41	38.002 39.178 39.453 40.269 40.368 40.454	3.73 7.54 5.24 18.23 3.10 2.11	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol 2,4a,8,8-Tetramethyl decahydro cyclopropa[d]naphthalene Olean-12-en-3-one 4,4,6a,6b,8a,11,11,14b-	C ₂₉ H ₅₀ O2 C ₂₈ H ₄₈ O C ₆ H ₁₂ O6 C ₂₀ H ₄₀ O C ₁₅ H ₂₆ C ₃₀ H ₄₈ O	430 400 180 296 206 424	
36 37 38 39 40	38.002 39.178 39.453 40.269 40.368	3.73 7.54 5.24 18.23 3.10	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol 2,4a,8,8-Tetramethyl decahydro cyclopropa[d]naphthalene Olean-12-en-3-one 4,4,6a,6b,8a,11,11,14b- Octamethyl1,4,4a,5,6,6a,6b,7,	C ₂₉ H ₅₀ O2 C ₂₈ H ₄₈ O C ₆ H ₁₂ O6 C ₂₀ H ₄₀ O C ₁₅ H ₂₆	430 400 180 296 206	
36 37 38 39 40 41 42	38.002 39.178 39.453 40.269 40.368 40.454 40.863	3.73 7.54 5.24 18.23 3.10 2.11 0.51	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol 2,4a,8,8-Tetramethyl decahydro cyclopropa[d]naphthalene Olean-12-en-3-one 4,4,6a,6b,8a,11,11,14b- Octamethyl1,4,4a,5,6,6a,6b,7, 8,8a,9,10,11,12,12a,	C29H50O2 C28H48O C6H12O6 C20H40O C15H26 C30H48O C30H48O	430 400 180 296 206 424 424	
36 37 38 39 40 41	38.002 39.178 39.453 40.269 40.368 40.454	3.73 7.54 5.24 18.23 3.10 2.11	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol 2,4a,8,8-Tetramethyl decahydro cyclopropa[d]naphthalene Olean-12-en-3-one 4,4,6a,6b,8a,11,11,14b- Octamethyl1,4,4a,5,6,6a,6b,7, 8,8a,9,10,11,12,12a, Stigmast-7-en-3-ol, (3. beta.,5.	C ₂₉ H ₅₀ O2 C ₂₈ H ₄₈ O C ₆ H ₁₂ O6 C ₂₀ H ₄₀ O C ₁₅ H ₂₆ C ₃₀ H ₄₈ O	430 400 180 296 206 424	
36 37 38 39 40 41 42 43	38.002 39.178 39.453 40.269 40.368 40.454 40.863 40.997	3.73 7.54 5.24 18.23 3.10 2.11 0.51 0.32	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol 2,4a,8,8-Tetramethyl decahydro cyclopropa[d]naphthalene Olean-12-en-3-one 4,4,6a,6b,8a,11,11,14b- Octamethyl1,4,4a,5,6,6a,6b,7, 8,8a,9,10,11,12,12a, Stigmast-7-en-3-ol, (3. beta.,5. alpha.,24S)-	C29H50O2 C28H48O C6H12O6 C20H40O C15H26 C30H48O C30H48O	430 400 180 296 206 424 424 424	
36 37 38 39 40 41 42	38.002 39.178 39.453 40.269 40.368 40.454 40.863	3.73 7.54 5.24 18.23 3.10 2.11 0.51	Vitamin E Ergost-5-en-3-ol, (3. beta.,24R)- D-Allose Phytol 2,4a,8,8-Tetramethyl decahydro cyclopropa[d]naphthalene Olean-12-en-3-one 4,4,6a,6b,8a,11,11,14b- Octamethyl1,4,4a,5,6,6a,6b,7, 8,8a,9,10,11,12,12a, Stigmast-7-en-3-ol, (3. beta.,5.	C29H50O2 C28H48O C6H12O6 C20H40O C15H26 C30H48O C30H48O	430 400 180 296 206 424 424	

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46	41.528	3.15	Pentadecanoic Acid	$C_{15}H_{30}O_2$	242	
47	41.734	1.14	. AlphaTocopheryl acetate	$C_{31}H_{52}O_3$	472	
48	42.090	2.04	. BetaAmyrin	C ₃₀ H ₅₀ O	426	
49	42.200	1.08	9,19-Cyclolanostan-3-ol, 24- methylene-, (3. beta.)-	C ₃₁ H ₅₂ O	440	
50	42.653	1.68	D: B-friedo-B': A'- Neogammacer-5-en-3-one	C ₃₀ H ₄₈ O	424	
51	42.866	7.50	9,12,15-Octadecatrienoic Acid,	$C_{18}H_{30}O_2$	278	
52	43.703	2.16	03027205002 Flavone 4'- OH,5-oh,7-di-O-Glucoside	$C_{27}H_{30}O_{15}$	594	
53	44.133	0.46	9,19-Cycloergost-24(28)-EN-3- ol, 4,14-dimethyl- ,(3.beta.,4.alpha., 1-Isopropenyl-4,5-	C ₃₀ H ₅₀ O	426	
54	44.281	0.99	dimethylbicyclo [4.3.0] nonan- 5-ylmethylphenyl sulfoxide	$C_{21}H_{30}OS$	330	
55	47.840	0.58	1-Eicosanol	$C_{20}H_{42}O$	298	
56	55.459	0.90 100.00	Phytol, acetate	C ₂₂ H ₄₂ O ₂	338	

Table 4: GC-MS report of methanolic extract of C. quadrangularis stem Variant III

Deel	R. Time	Area9/		Molecular	Molecular
Peak	R. Time	Area%	Name of the compounds	formula	weight
1	24.445	0.62	n-Tetracosanol-1	C ₂₄ H ₅₀ O	354
2	24.517	0.21	Stigmasterol	$C_{29}H_{48}O$	412
3	26.398	0.27	Cyclohexane, 1,1'- tetradecylidenebis-	C ₂₆ H ₅₀	362
4	28.943	1.08	cis-9-Hexadecenal	C ₁₆ H ₃₀ O	238
5	30.243	0.50	Heptacosyl acetate	C ₂₉ H ₅₈ O ₂	438
6	31.683	1.11	Squalene	C ₃₀ H ₅₀	410
7	33.064	2.00	Beta-Sitosterol	C ₂₉ H ₅₀ O	414
8	33.469	0.40	Eicosanal-	C ₂₀ H ₄₂ O	298
9	34.544	2.76	Octanoic acid, 2-di methylaminoethyl ester	C ₁₂ H ₂₅ NO ₂	215
10	36.341	2.83	Celidoniol, deoxy-	C ₂₉ H ₆₀	408
11	38.532	2.78	5-Phenyl-Pentanoic Acid, Ethyl Ester	C17H24O5	308
12	40.096	2.13	Tetracontane	C40H82	562
13	40.226	0.83	9,19-Cycloergost-24(28)-EN- 3-ol, 4,14-dimethyl- ,(3.beta.,4.alpha.,	C ₃₀ H ₅₀ O	426
14	41.182	2.51	Hexatriacontane	C36H74	506
15	42.109	1.58	Fumaric acid, 2-di methyl aminoethylnonyl ester	C ₁₇ H ₃₁ NO ₄	313
16	42.205	0.27	1-Docosanol, acetate	C24H48O2	368
17	42.513	0.41	Phytol, acetate	$C_{22}H_{42}O_2$	338
18	42.583	0.72	gammaTocopherol	C ₂₈ H ₄₈ O ₂	416
19	43.112	1.81	Tetratetracontane	$C_{44}H_{90}$	618



		100.00			
35	50.920	0.56	Stigmastane-3,6-dione, (5. alpha.)-	C29H48O2	428
34	49.601	2.17	03027205002 Flavone 4'- OH,5-OH,7-Di-O-Glucosid	C ₂₇ H ₃₀ O ₁₅	594
33	48.719	0.76	24-Norursa-3,12-diene	C ₂₉ H ₄₆	394
32	48.363	1.28	9,19-Cyclo-9. beta lanostane-3. beta.,25-diol	$C_{30}H_{52}O_2$	444
31	47.853	6.82	3-O-Methyl-d-glucose	$C_7H_{14}O_6$	194
30	47.285	3.71	Phytol	C ₂₀ H ₄₀ O	296
29	46.730	8.10	9,12,15-Octadecatrienoic Acid,	C ₁₈ H ₃₀ O ₂	278
28	46.537	3.11	4,4,6A,6B,8A,11,11,14B- Octamethyl-1,4,4A,5,6,6A,6	C ₃₀ H ₄₈ O	424
27	46.005	23.24	D-Allose	$C_6H_{12}O_6$	180
26	45.866	16.27	Hexadecanoic acid	C ₁₆ H ₃₂ O ₂	256
25	45.529	0.84	n-Tetratetracontane	C44H90	618
24	45.039	3.09	. BetaAmyrin	C ₃₀ H ₅₀ O	426
23	44.810	1.32	Beta.,24R)- Henicosanal	C21H42O	310
22	44.697	3.14	Ergost-5-en-3-ol, (3.	C ₂₈ H ₄₈ O	400
21	44.374	0.41	Octacosyl acetate	$C_{30}H_{60}O_2$	452
20	44.245	0.37	Alpha-d-galactopyranose, 6- O-(2,3,5-TRI-O-AC	$C_{23}H_{34}O_{13}$	518

FT-IR

FT-IR range was utilized to distinguish the utilitarian gathering of the dynamic segments in view of the peak an incentive in the point of infrared radiation. The FT-IR analysis results showed that the methanolic stem extract of *C. quadrangularis* variant I having the presence of Alcohol, Aldehyde, Iso cyanides, Alkane, Primary alcohol, Chloro constituent, which shows major peaks at 3420.91 cm⁻¹, 2924.27 cm⁻¹, 2846.85 cm⁻¹, 2076.74 cm⁻¹, 1647.04 cm⁻¹, 1454.75 cm⁻¹, 1409.96 cm⁻¹, 1053.44 cm⁻¹, 1032.56 cm⁻¹, 1017.56 cm⁻¹, 718.75 cm⁻¹ and 666.26 cm⁻¹ respectively (Figure 7 & Table 5). The FT-IR analysis of methanolic stem extract of *C.*

quadrangularis variant II, the presence of Alcohol, Aldehyde, Iso cynides, Nirite, Alkane, Carboxylic acid, Tertiary alcohol, Primary alcohol and Chloro compound. Constituent which shows major peaks at 3434.81 cm⁻¹, 2867.19 cm⁻¹, 2077.67 cm⁻¹, 1640.48 cm⁻¹, 1454.51 cm⁻¹, 1400.70 cm⁻¹, 1108.01 cm⁻¹, 1054.30 cm⁻¹ and 678.41 cm⁻¹ respectively (Figure 8 & Table 6). The FT-IR analysis methanolic stem extract of *C. quadrangularis* variant III showed Alcohol, Aldehyde, Iso cyanides, Alkyl compound, Alkane, Primary alcohol, Chloro compound. Constituent which shows high peaks at 3434.32 cm⁻¹, 2842.29 cm⁻¹, 2519.69 cm⁻¹, 1646.67 cm⁻¹,1456.31 cm⁻¹, 1017.38 cm⁻¹ and 574.13 cm⁻¹ (Figure 9 & Table 7).

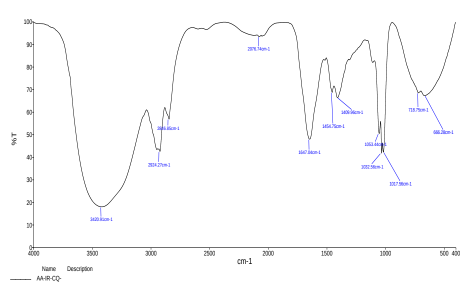


Figure 7: FT-IR Spectrum of methanolic extract of *C. quadrangularis* stem variant I

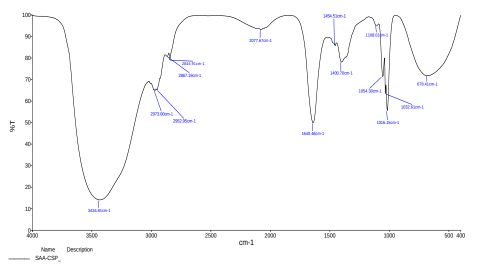


Figure 8: FT-IR Spectrum of methanolic extract of C. quadrangularis Stem variant II

S. No	Wave No.	Molecular Motion	Functional group	Absorption intensity
1	3420.91cm ⁻¹	O-H	Alcohol	Strong
2	2924.27 cm ⁻¹	O-H	Alcohol	Medium
3	2846.85 cm ⁻¹	C=O	Aldehyde	Weak
4	2076.74 cm ⁻¹	C=N	lso cyanides	Medium
5	1647.04 cm ⁻¹	O-NO	Nirite	Strong
6	1454.75 cm ⁻¹	C-H	Alkane	Medium
7	1409.96 cm ⁻¹	C-H	Alkane	Medium
8	1053.44 cm ⁻¹	C-0	Primary alcohol	Strong
9	1032.56 cm ⁻¹	C-0	Primary alcohol	Strong
10	1017.56 cm ⁻¹	C-0	Primary alcohol	Strong
11	718.75 cm ⁻¹	C-Cl	Chloro compound	Strong
12	666.26 cm ⁻¹	C-Cl	Chloro compound	Strong

Table 5: FT-IR absorption and functional groups of stem extract of *C. quadrangularis* variant I



S. No	Wave No.	Molecular Motion	Functional group	Absorption intensity
1	3434.81 cm ⁻¹	O-H	Alcohol	Medium
2	2973.00 cm ⁻¹	O-H	Alcohol	Medium
3	2952.95 cm ⁻¹	O-H	Alcohol	Medium
4	2867.19 cm ⁻¹	C=0	Aldehyde	Weak
5	2844.91 cm ⁻¹	C=0	Aldehyde	Weak
6	2077.67 cm ⁻¹	C=N	lso cynides	Medium
7	1640.48 cm⁻¹	O-NO	Nirite	Strong
8	1454.51 cm ⁻¹	C-H	Alkane	Medium
9	1400.70 cm ⁻¹	C-0	Carboxylic acid	Weak
10	1108.01 cm ⁻¹	C-0	Tertiary alcohol	Strong
11	1054.30 cm ⁻¹	C-0	Primary alcohol	Strong
12	1032.61 cm ⁻¹	C-0	Primary alcohol	Strong
13	1016.15 cm ⁻¹	C-0	Primary alcohol	Strong
14	678.41 cm ⁻¹	C-Cl	Chloro compound	Strong

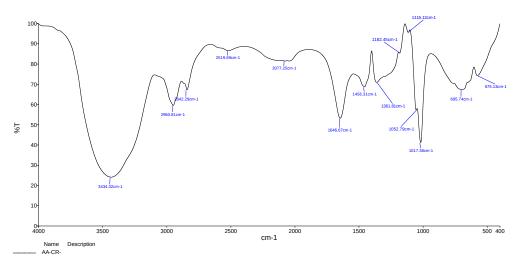


Figure 9: FT-IR Spectrum of methanolic Stem extract of *C. quadrangularis* variant III

S. No	Wave No.	Molecular Motion	Functional group	Absorption intensity
1	3434.32 cm ⁻¹	0-Н	Alcohol	Strong
2	2950.81 cm ⁻¹	O-H	Alcohol	Medium
3	2842.29 cm ⁻¹	C=0	Aldehyde	Weak
4	2519.69 cm ⁻¹	C=N	Iso cyanides	Medium
5	2077.25 cm ⁻¹	C=N	Iso cyanides	Medium
6	1646.67 cm ⁻¹	C=N	Alkyl compound	Strong
7	1456.31 cm ⁻¹	C-H	Alkane	Medium
8	1361.81 cm ⁻¹	C-H	Alkane	Medium
9	1182.45 cm ⁻¹	C-0	Primary alcohol	Strong
10	1115.12 cm ⁻¹	C-0	Primary alcohol	Strong
11	1052.79 cm ⁻¹	C-0	Primary alcohol	Strong
12	1017.38 cm ⁻¹	C-0	Primary alcohol	Strong
13	695.74 cm ⁻¹	C-Cl	Chloro compound	Strong
14	574.13 cm ⁻¹	C-Cl	Chloro compound	Strong

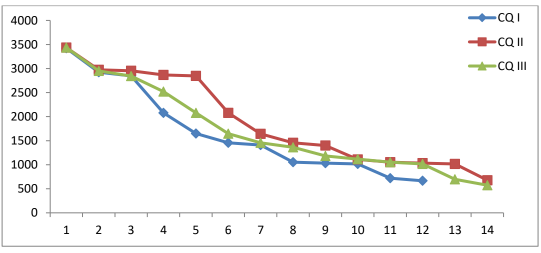
Table 7: FT-IR absorption and functional groups of stem extract of C. quadrangularis variant III

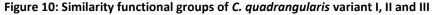


Comparison of phytochemical compounds from *C. quadrangularis* all three-variant based on GC-MS and their pharmacological activities

Based on the GC-MS and FT-IR analyses were identified phytochemical compounds and functional groups which present in the methanolic Stem extract of C. quadrangularis Variant I, II and III. All the three variant of C. quadrangularis contain 3- o- methyl- d- glucose, D-Allose, Phytol, 9, 12, 15 - octadecatrienoic acid. These four compounds are having Anti-tubercular activity, allergic disorders and anti-microbial activities, joint disorders, wounds healing activities. Pentadeconic acid and vitamin E were present in C. quadrangularis variant I and II. Pharmacology activities of this compounds such as Anti-cancer drug, anti-asthmatics and anti-abortive and sexual disorders. C. quadrangularis variant II and III contains similar phyto-components such as hexadeconic acid, ergost – 5-en- 3- ol and Beta- amyrin. Hexadeconic acid was major active compound of the variant II and III. These three compounds having bone healing activity

and wound healing activity. Beta - sitosterol only one compound found in C. quadrangularis variant I and III. This compound has been used for anti-cancer, bone and wound healing activities. Based on the phytochemical analysis similar phyto-components present in the all three variants and having bone and wound healing activities. Pharmacological activities of С. quadrangularis Variant I and II has been report from earlier [11]. Presence of 3- o- methyl- d- glucose, D-Allose, Phytol, 9, 12, 15 - octadecatrienoic acid, hexadeconic acid, ergost - 5-en- 3- ol and Beta- amyrin. Hexadeconic acid and Beta- sitosterol of Methanolic Stem extract of C. quadrangularis Variant III also used as pharmacology activities. Based on the FT-IR analysis similar functional group were fund in all the three variant of C. quadrangularis. Such as alcohol, aldehyde, iso cyanide, alkane and Cholo compound. Nirite functional groups of compounds present in C. quadrangularis variant I and II. Carboxylic acid fund in C. quadrangularis variant II. (Figure 10).





CONCLUSION

Based on the morphology of the stem character of C. quadrangularis all the three variants showed distinct shaped stems. Based on the GC-MS analyses of C. quadrangularis variants showed similar major compounds like viz., 3- o- methyl- d- glucose, D- Allose, Phytol, 9, 12, 15 - octadecatrienoic acid. These compounds were used for anti-cancer, joint disorders, wounds healing activities and some other pharmacological activities. FT-IR analyses also showed similar functional group of compounds related with same medicinal properties.

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Received:04.05.18, Accepted: 07.06.18, Published:01.07.2018

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